



DATA PACK - KIMTECH PURE* A5 Sterile Cleanroom Apparel **Contents**

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KIMTECH PURE* A5 Sterile Cleanroom Coveralls.

The KIMTECH PURE* A5 Sterile Cleanroom Coveralls simplify aseptic gowning to save time and reduce the risk of contamination.



CLEAN-DON* blue line on the inside indicates correct place to grasp for aseptic gowning.



Telescope fold and snaps prevent legs and arms falling to the floor during gowning.



Vacuum Packed and double bagged for added sterility assurance.

Reduces space so more products fit into gowning area storage.









STERILE R

PERSONAL & PROCESS PROTECTION:

- Suitable for EU GMP ISO 5 Grade A cleanrooms
- Gamma Irradiated, Sterility Assurance Level 10-6
- Low Lint fabric, Helmke Drum test Category I
- Bacterial Filtration Efficiency 96%
- Manufactured and Packed in ISO 5 cleanroom
- Vacuum packed for sterility assurance, saves space
- Certified PPE Cat III directive 89/686/EEC
- Type 6 limited chemical splash protection
- Type 5 particle protection
- · Breathable barrier combines comfort and protection

DESIGN FEATURES:

- Unhooded coverall available in size S to 4XL
- Blue Indicator line to avoid touching outside
- Presented unzipped with inside-out fold
- Arms and legs telescope folded with snaps
- Waist and back panel elastics
- Elastic cuffs and thumb loops
- High strength, bound and triple stitched seams
- High performance SMS breathable barrier
- Individually vacuum packed and double bagged.
- 25 items per double case liner.

Applications:

EU GMP ISO 5 Grade A

Aseptic Processing

Parenteral
Drug
Manufacturing

Biotechnology

Pharmaceutical Compounding

Ophthalmic product manufacturing



KIMTECH PURE* A5 Sterile Integrated Hood & Mask.

The KIMTECH PURE* A5 Sterile Integrated Hood & Mask simplifies aseptic gowning to save time and reduce the risk of contamination.



CLEAN-DON* blue line on the inside indicates correct place to grasp for aseptic gowning. Saves time and reduces contamination risk.



No gaps between hood and mask ensures exhaled air is filtered.

Elastics provide a comfortable, secure and universal fit.



Vacuum packed for added sterility assurance. Reduces space so more products fit into gowning area storage.









PERSONAL & PROCESS PROTECTION:

- Suitable for EU GMP ISO 5 Grade A cleanrooms
- Gamma Irradiated, Sterility Assurance Level 10-6
- Low Lint fabric, Helmke Drum test Category I
- Bacterial Filtration Efficiency 96%
- Manufactured and Packed in ISO 5 cleanroom
- Vacuum packed for sterility assurance, saves space
- Certified PPE Cat III directive 89/686/EEC
- Type 6 Pb limited chemical splash protection
- Breathable barrier combines comfort and protection

DESIGN FEATURES:

- Integrated hood and mask
- · Stretch fit elastics for universal size
- Blue Indicator line to avoid touching outside
- Pull down ties on back to assist gowning
- Presented inside-out fold to avoid contamination
- High strength, bound and triple stitched seams
- High performance SMS with cloth-like feel
- Individually vacuum packed
- 25 items per double-bag, 3 double bags per case

Applications:

EU GMP ISO 5 Grade A Aseptic Processing Parenteral
Drug
Manufacturing

Biotechnology

Pharmaceutical Compounding

Ophthalmic product manufacturing



KIMTECH PURE* A5 Sterile Cleanroom Boot Covers.

The KIMTECH PURE* A5 Sterile Boot Covers improves worker safety and comfort, while reducing the risk of contamination.



New Version Ideal for wet applications. Wrap-around Vinyl foot is seamed above the floor line to reduce risk of liquid penetration and trip hazards. Standard vinyl sole also available.



3 sizes for better safety and comfort. Better fit and comfort to improve worker safety by reducing risk of trip hazards



Vacuum packed for added sterility assurance. Reduces space so more products fit into gowning area storage.







PROCESS PROTECTION:

- Suitable for FU GMP ISO 5 Grade A cleanrooms.
- Gamma Irradiated, Sterility Assurance Level 10⁻⁶
- Low Lint fabric, Helmke Drum test Category I
- Bacterial Filtration Efficiency 96%
- Manufactured and Packed in ISO 5 cleanroom
- Vacuum packed for sterility assurance, saves space
- CE 89/686/EEC Category I PPE simple design
- Breathable barrier combines comfort and protection

DESIGN FEATURES:

- New Vinvl foot version available in 3 sizes.
- Standard Vinyl sole version available in 3 sizes.
- Elastic opening presented inside-out folded.
- Two extra-long ties.
- SMS material cloth-like feel keeps gloves in place
- High strength, bound and triple stitched seams
- Individually vacuum packed pairs
- 25 pairs per double-bag, 4 double bags per case

Applications:

EU GMP ISO 5 Grade A Aseptic

Parenteral | Drua Manufacturing

Biotechnology

Pharmaceutical Compounding

Ophthalmic product manufacturing

KIMTECH PURE* A5 Sterile Cleanroom Sleeves.

The KIMTECH PURE* A5 Sterile Cleanroom Sleeves simplify aseptic gowning to save time and reduce the risk of contamination.



CLEAN-DON* blue line on the inside indicates correct place to grasp for aseptic gowning.

Thumb-loops and material's cloth-like feel keep sleeves and gloves in place.



Telescoped fold makes aseptic preventing gowning easier, contact with the outside surface to reduce contamination risk.



Vacuum packed for added sterility assurance. Reduces space so more products fit into gowning area storage.







PROCESS PROTECTION:

- Suitable for FU GMP ISO 5 Grade A cleanrooms.
- Gamma Irradiated, Sterility Assurance Level 10⁻⁶
- Low Lint fabric, Helmke Drum test Category I
- Bacterial Filtration Efficiency 96%
- Manufactured and Packed in ISO 5 cleanroom
- Vacuum packed for sterility assurance, saves space
- CE 89/686/EEC Category I PPE simple design
- Breathable barrier combines comfort and protection

DESIGN FEATURES:

- 45cm long sleeve protectors, pair packed.
- Blue Indicator line to avoid touching outside
- Telescope inside-out fold for easy gowning
- Elastic cuffs and thumb loops
- SMS material cloth-like feel keeps gloves in place
- High strength, bound and triple stitched seams
- Individually vacuum packed pairs
- 25 pairs per double-bag, 4 double bags per case

Applications:

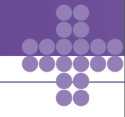
EU GMP ISO 5 Grade A Aseptic

Parenteral | Drug Manufacturing

Biotechnology

Pharmaceutical Compounding

Ophthalmic product manufacturing



KIMTECH PURE* A5 Sterile Cleanroom Apparel **Technical Data**

Quality Standards

- Suitable for ISO 5 Grade A Sterile Cleanrooms
- Manufactured in ISO9001 and ISO13485 certified facility
- Sterility Assurance Level 10⁻⁶ and Helmke Drum test Category I
- Sterilization in accordance with EN556-1, EN ISO11737-1,-2, EN ISO 11140-1

Fabric Tests	Test Method	Result
Particle shedding test (Helmke Drum)	IEST-RP-CC003.3	Category I
Bacterial Filtration Efficiency (3.0 µm) - SMS material in A5 fabric	ASTM F2100	96%
Bacterial Filtration Efficiency (3.0 µm) - Mask in iHAM	ASTM F2100	94%
Particle filtration Efficiency (0.5 $\mu m)$ - SMS material A5 fabric	ASTM F2299	94%
Particle Filtration Efficiency (0.5 µm) - Mask in iHAM	ASTM F2299	93%
Abrasion resistance	EN 530 Method 2	Class 3 of 3
Flex cracking resistance	ISO 7854 Method B	Class 5 of 6
Trapezoidal tear resistance	ISO 9073-4	Class 2 of 3
Puncture resistance	EN 863	Class 1 of 3
Tensile strength	EN ISO 13934-1	Class 1 of 3
Resistance to ignition	EN 13274-4 Method 3	Pass
Seam strength	EN ISO 13935-2	Class 3 of 3

EN ISO 6350:2005 — Resistance of penetration by liquids/chemicals.	CAS number	Penetration Class	Penetration %	Repellency Class	Repellency %
1-butanol 100%	71-36-3	3	<1%	1	>80%
Ethanol 70%	64-17-5	3	<1%	2	>90%
Ethylene glycol 100%	107-21-1	3	<1%	2	>90%
Incidin Plus 100%	N/A	3	<1%	2	>90%
Isopropyl Alcohol 70%	67-63-0	2	<5%	1	>80%
Sekusept plus	N.A	3	<1%	2	>90%
Sodium Hydroxide 10%	1310-73-2	3	<1%	3	>95%
Sulphuric Acid 30%	7664-93-9	3	<1%	2	>90%

KIMTECH PURE* A5 Sterile Cleanroom Apparel

Description	Code.
Coveralls, 25 per Case	88800 (S), 88801 (M), 88802 (L), 88803 (XL), 88804 (2XL), 88805 (3XL), 88806 (4XL)
Boots Vinyl Foot, 100 pairs per Case	12922 (S), 88808 (Universal), 12920 (XL)
Boots Vinyl Sole, 100 pairs per Case	31683 (S), 31696 (Universal), 31697 (XL)

Description	Code.
Integrated Hood & Mask, 75 per Case	36072
Hood with Ties, 100 per Case	25797 / 88807
Sterile Sleeves, 100 pairs per Case	36077
Sterile Sleeves, 100 pairs per Case	36077





KIMTECH PURE* A5 Sterile Cleanroom Apparel with CLEAN-DON* Technology

Gowning Procedure BEFORE GOWNING

Step 1: (Pre-Entry) Don Hair Net and Shoe Covers after removing all jewelry and cosmetics.

Step 2: (Gowning) Wash hands and gown first pair of sterile gloves. Sanitize gloves after gowning each article if required.

Step 3: Apply mask and hood assuring a snug fit.

Step 4: Open vacuumed-packed apparel. Tear at notched edge.



Step 5: Grasp the blue line. Located on the inside middle back.

Step 6: Gently unfold coverall utilizing blue indicator line. Arms and legs are pre-drawn and snapped in place. Garment is already folded inside-out and unzipped.

BEGIN GOWNING

Snaps allow gathered-up arms and legs to expand during gowning



Step 7: Hold garment at waist



Step 8: Put one leg in and point toe through opening until snap releases



Step 9: Do the same with the other leg



Step 10: Insert one arm and extend until snap releases









Final Step: Cross legs and zip up coverall



Add boot covers



Complete gowning by adding goggles and a second pair of sterile gloves.



KIMTECH PURE* A5 Integrated Hood and Mask Donning Technique



Step 1

Select your integrated hood and mask; inspect package for vacuum seal; tear along notches toward the product to open



Step 2

Pull out integrated hood and unfold to locate the blue signal indicator line



Step 3

Center your chin just above the blue line then roll over your forehead.



Step 4

Grab ties with both hands then extend arms out in a circular motion to position under armpits.



Step 5

Pull ties across chest and secure as needed



Step 6

Gently adjust front flap to comfortable position; pinch mask wire over bridge of nose to ensure proper fit.





KIMTECH PURE* A5 Sterile Boot Cover with Grasp Ties Donning Technique



Step 1

Select your boot cover size; inspect package for vacuum seal; tear along notches toward the product to open; flex package to separate boot covers



Step 2

Pull out one boot cover by the cuff



Step 3

Grasp ties with one hand and make sure they don't touch the floor



Step 4

Open up boot cover; point toe towards opening and pull up over calf while holding the ties



Step 5

Hold up foot or rest on bench. Grasp ties at front of boot cover and wrap over cusp of shoe/boot; then wrap ties behind ankle and tie in front.



Step 6

Place donned boot on the clean side



Step 7

Pull the remaining boot cover out of package by the cuff and repeat Steps 3, 4 and 5. Proceed with final gowning steps







A5 Sterile Cleanroom Coveralls 88800 (S), 88801 (M), 88802 (L), 88803 (XL) 88804 (2XL), 88806 (3XL), 88806 (4XL)

A5 Sterile Cleanroom Sleeves 36072 — one size fits all

Case Label





Item Label







A5 Sterile Cleanroom Hoods with ties EU 88807(01) / US 25797(00)

A5 Sterile Integrated Hood & Mask 36072 – one size fits all

Case Label





Item Label



Please note:

From June 2013 production, 88807(01) will be dual coded to 25797 (US) / 88807 (EU).

This product is identical in design and specification.

Our European business will continue to refer to this product as 88807 in all price lists, literature and ordering systems.





A5 Sterile Boot Covers with Vinyl Sole 12922 (S), 88808 (Universal), 12920 (XL)

A5 Sterile Boot Covers with Vinyl Foot 31683 (S), 31696 (Universal), 31697 (XL)

Case Label













Certificate of Conformance KIMTECH PURE* A5 Cleanroom Sterile Apparel

Product: KIMTECH PURE* A5 Cleanroom Sterile Apparel

Code	Description	Size	Packaging
88800	Coverall	S	25/case
88801	Coverall	М	25/case
88802	Coverall	L	25/case
88803	Coverall	XL	25/case
88804	Coverall	2XL	25/case
88805	Coverall	3XL	25/case
88806	Coverall	4XL	25/case
12917	Coverall	5XL	25/case
12914	Coverall	6XL	25/case
88807	Hood, no ties	One-Size	100/case
25797	Hood, with ties	One-Size	100/case
88808	Universal Boots	One-Size	100 pr/case
12922	Boots	S/M	100 pr/case
12920	Boots	XL/2XL	100 pr/case
31683	Boots, with edge vinyl	S/M	100 pr/case
31696	Boots, with edge vinyl	One-Size	100 pr/case
31697	Boots, with edge vinyl	XL/2XL	100 pr case

Lot number: XN213701X

This document certifies that the lots listed above conform to Kimberly-Clark's internal specifications for product quality. Kimberly-Clark uses a system of in-process and lot inspections to assure conformance to specifications.

Characteristic	Specification Target
Particle Count (Helmke Drum, IEST-RP-CC003.3)	Category I ¹
Sterility Assurance Level (ANSI/AAMI/ISO 11137)	10 ⁻⁶

A Certificate of Irradiation is also available to assure that the product has received the specified radiation dosage.

	Tiangh Wan			
Verified by:		Date:	May 29, 2012	
® Registered Tradema	rk and * Trademark of Kimberly-Clark \	Worldwide Inc © 2006	S KCWW All Rights Reserve	d

¹Product is tested prior to sterilization Customer Service: 800-255-6401

辐照证明书

CERTIFICATE OF IRRADIATION



深圳市金鹏源辐照技术有限公司

SHENZHEN JPY 10N-TECH. CO., LTD. 地址 (Add): 深圳市罗湖区市心东盛路68号 No.68 Dongsheng Rd, Buxin, Lobou, Shenzhen 18191 China 电 话(TEL):+86(0)755 25177231, 25177137 传 真(FAX):+86(0)755 25516854, 邮政编码 (FC.): 518019 http://www.jpy.com.cn

R/SZJPY(B)-7515 AC/01

证书编号:

Certificate No.:

GM2012071502

分合同编号:

以下空白

Sub-contract:

GM12070127

产品名称 Article description	包装规格 (厘米) Carton Size(cm)	产品批号 Article Lot No.	数量(箱) Carton Qty(cs)	毛重(公斤) G.W(kgs)
Universal Boots:8880	50*30*37	XN218101X	*180*	1620
Sleeve: 36077	50*30*37	XN218801X	*62*	334.8
Coveral1: 88802	50*30*37	XN213701X	*22*	158.4
Coverall: 88805	50*30*42	XN213701X	*75*	645
Universal Boots:8880	50*30*37	XN218801X	*40*	360

这批产品已经过伽玛射线辐照。

THE PRODUCTS HAVE BEEN IRRADIATED BY GAMMARAY

THE I RODGETS HAVE BEEN IKKA	DIAILD DI GAMMAKAI.
加工编号 Process Code No.	P1207153
辐照结束的日期 Irradiation lot finish date	2012-07-15
客户要求最低吸收剂量: Minimum specified dose: 32.5 kGy	最低剂量监测区吸收剂量: Minimum inspection area dose: 34.0 kGy
客户要求最高吸收剂量: Maximum specified dose: 50 kGy	最高剂量监测区吸收剂量: Maximum inspection area dose: 44.6 kGy
检验员: Inspector: /以及 2012.7.15	深圳市金鹏源辐照技术有限公司
审核: Approved: 文 知道	光明分公司 SHENZHEN JPY ION-TECH. CO., LTD.
日期: Date: 2012 - 7、15	GUANGMING BRANCH



A Bivision of Sterigenics International, Inc.

Protocol Number	Revision	Issue Date	Brief Description of Revision
797100277-P	0	April 12, 2010	
March 1964	1		
	2		

SteriPro Consul Sponsor Name	tant: Zabrina Tumaitis	Sponsor Contact:
Kimberly-Clark	and Address.	'
1400 Holcomb B	Bridge Road	Theresa McCoy
Roswell, GA 30	076	
Prepared By:	Zabrina Tumaitis Consultant, SteriPro Consulting	Date
Reviewed By:		Date
	Julie Arinaga Quality Assurance Manager, SteriPro Labs	
Approved By:		Date
	Niki Fidopiastis Director, SteriPro Consulting	Date
Approved By:	Theresa McCoy	April 14,20



A Division of Sterigenics International, Inc.

Protocol Number	Revision	Issue Date	Brief Description of Revision
797100277-P	0	April 12, 2010	
	1		
	2		

Protocol Title		
VD-Ma	x32.5 Radiation Validation Cleanroom Sterile	
	tant: Zabrina Tumaitis	
Sponsor Name : Kimberly-Clark	and Address:	Sponsor Contact:
1400 Holcomb E Roswell, GA 300		Theresa McCoy
Prepared By:	Zabrina Tumaitis Consultant, SteriPro Consulting	Date
Reviewed By:	Julie Arinaga Quality Assurance Manager, SteriPro Labs	Date
	Niki Fidopias	Digitally signed by Niki Fidopiastis DN: cn=Niki Fidopiastis, o=Director of Consulting, ou=SteriPro, email=nfidopiastis@sterigenics.com, c=U5 Date: 2010.04.14 07:34:02 -07'00'
Approved By:	Niki Fidopiastis Director, SteriPro Consulting	Date
Approved By:	Theresa McCoy Kimberly-Clark	Date
period of five (aintain copies of all Protocols, Final Rep 5) years from the report issue date. Afton hese materials. Confidential	ports, and supporting documentation for a retender this period, SteriPro will contact the sponsor



A Division of
Sterigenics International, Inc.

Protocol Number	Revision	Issue Date	Brief Description of Revision
797100277-P	0	April 12, 2010	
	1		
	2		

Protocol Title VD-Ma	x32.5 Radiation Validation for K Cleanroom Sterile Appa	
SteriPro Consul Sponsor Name	tant: Zabrina Tumaitis	Sponsor Contact:
Kimberly-Clark 1400 Holcomb E Roswell, GA 300	Bridge Road	Theresa McCoy
Prepared By:	Zabrina Tumaitis Consultant, SteriPro Consulting	12Aprilu Date
Reviewed By:	Julie Arinaga Quality Assurance Manager, SteriPro Labs	
Approved By:	Niki Fidopiastis Director, SteriPro Consulting	Date
Approved By:	Theresa McCoy Kimberly-Clark	Date

Confidential

disposition of these materials.



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Terms and Conditions of Study

Any and all reports and letters issued by SteriPro Consulting or SteriPro Laboratories, divisions of Sterigenics U.S., LLC (collectively referred to as "Sterigenics") are for the sole and exclusive use of Sterigenics and the sponsor to whom they are addressed. Sterigenics strictly prohibits any other use or reproduction of any reports or portions thereof, quotations from any such reports or use of Sterigenics' name without Sterigenics' prior written consent. Sponsor acknowledges and shall be deemed to have agreed, upon its use of Sterigenics' services that the significance of any data is subject to the adequacy and representative character of the samples tendered for testing. Sterigenics warrants only that all tests are performed in accordance with established laboratory procedures and standards. Sterigenics makes no other warranties or representations of any kind, express or implied, including but not limited to anything regarding the adequacy of the samples tendered for testing for any specific use or application, such determination being the sole responsibility of the sponsor. Sponsor acknowledges and shall be deemed to have agreed, upon its use of Sterigenics' services, that Sterigenics' liability for any loss or damage resulting from Sterigenics' acts or omissions shall in no event exceed the cost of tests performed. In no event shall Sterigenics be liable for any special, indirect, incidental, consequential, punitive or other similar damages, including but not limited to damages arising from death, bodily injury, property damage (other than as set forth herein), loss of profits or revenue or loss of use of any samples or products.



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1. Scope

This study will be performed to establish a 32.5-kGy minimum dose and validate the effectiveness of Gamma Radiation sterilization of the KIMTECH PURE* A5 Cleanroom Sterile Apparel. Please refer to the Product Family Members Table below for all products included in this sterilization family. This protocol describes procedures for validation of radiation sterilization of medical products in accordance with AAMI TIR 33: Sterilization of health care products – Radiation – Substantiation of a selected sterilization dose – Method VD-Max.

Product Family Members				
Description				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Coveralls - Size S				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Coveralls - Size M				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Coveralls - Size L				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Coveralls - Size XL				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Coveralls - Size 2XL				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Coveralls - Size 3XL				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Coveralls - Size 4XL				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Coveralls - Size 5XL				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Coveralls - Size 6XL				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Hoods - One size fits all				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Boots - Size S/M				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Boots - Size L				
Kimtech Pure* A5 Cleanroom Sterile Apparel - Boots - XL/2XL				

2. Objectives

- 2.1. Preliminary bioburden data has determined the Kimtech Pure* A5 Cleanroom Sterile Apparel Coveralls Size 6XL as the Dose Setting Device for this sterilization family.
- 2.2. The Dose Setting Device for the product family will be the test sample for this validation. Presterilization bioburden level will be determined to establish the appropriate verification dose.
- 2.3. Recommendation for a routine minimum sterilization dose will be based on evaluation of microbial survivors following exposure of products to the verification dose. The minimum dose of 32.5 kGy will be designed to provide a Sterility Assurance Level (SAL) of 10⁻⁶ or not more than one non-sterile unit for each one million units sterilized at that dose level.
- 2.4. This study is intended to support release of products for which exposure to the minimum required dose can be demonstrated by the use of calibrated dosimeters and without post-exposure sterility testing of each lot.



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3. Rationale

- 3.1. The VD-Max validation is utilized to determine the minimum effective processing dose for radiation sterilization. The validation is based on the concept that the product bioburden for the overall average or a single lot average (if greater than twice the overall average) bioburden will be less than 100,000 colony forming units.
- 3.2. The VD-Max procedure analyzes the number and resistance of the bioburden on the product. Three independent lots of product are tested for bioburden levels, and the bioburden recovery efficiency (percent recovery) is determined. The bioburden recovery efficiency and the sample item portion (percent of unit tested) are used to calculate the theoretical device bioburden on each test unit. The final bioburden estimate is used to determine the verification dose from the radiation dose tables specified in AAMI TIR 33.
- 3.3. The verification dose (sub-lethal) provides a SAL of 10⁻¹. A verification experiment must be performed to verify that the product bioburden resistance is less than or equal to the standard distribution of resistances used in the AAMI radiation guidelines.
- 3.4. Following completion of the study, a final report shall be generated by SteriPro, which shall be signed and approved by designated individuals from SteriPro, and Kimberly-Clark. This report shall be submitted to Kimberly-Clark and SteriPro will retain a copy as specified in the SteriPro procedure for record retention.

4. Terms and Definitions

Refer to ANSI/AAMI/ISO 11137:2006, AAMI TIR 33, and ANSI/AAMI/ISO 11737:2006 for definitions of terminology.

5. Responsibilities

5.1. Contract Irradiator

Sterigenics, Corona or Charlotte will be responsible for all equipment, process qualification, dosimeter calibration, and all documentation involving the ExCell irradiator's validation. A certificate of processing will be provided for the product in this study. All other Sterigenics records that are specific to this study will be available to the sponsor upon request or audit.

5.2. Contract Laboratory

SteriPro Labs, Itasca will perform the necessary microbiological testing for this validation and will provide all necessary documentation for the Final Report. DNV Certification, certificate numbers 2005-OSL-AQ-7662 and 2005-OSL-AQ-0212 certify SteriPro Labs to ISO 9001 and ISO 13485.

6. Product Specifications

6.1. Product Sample Amounts

Forty-eight (48) finished, routine product samples will be submitted in standard, final packaging format for all testing. These units shall be pulled after all steps of production except sterilization and be produced under current good manufacturing practice conditions. From the 48 product samples, there shall be three independent lots of ten samples each for bioburden determination. The 18 remaining samples may come from one or more production lots.



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6.2. Sample Item Portion (SIP)

The SIP used for all testing will be 0.10. The SIP that will be tested as the test sample will consist of different parts of the coverall for a total SIP of 0.10.

6.3. Sample Preparation

Additional sample prep is needed prior to sterilization in order to facilitate testing.

7. Bioburden Study Method

- 7.1. Recovery Determination
 - 7.1.1. Efficiency of Recovery Factor (ERF) will be determined using a specific bioburden recovery test procedure, which will be performed on a minimum of five samples taken from one or more production lots. This test is performed to determine the percentage of microorganisms that can be recovered from a product.
 - 7.1.2.Averaging the recovery obtained from the samples tested will derive the recovery factor. The bioburden test results for each lot will be adjusted by applying the ERF, in order to obtain a theoretical bioburden estimate. This theoretical bioburden estimate is a more accurate representation of the actual number of microorganisms on the product.
- 7.2. Recovery Test Method

The bioburden recovery test method for the product will be performed using the repetitive extraction method or inoculated recovery method. The extraction method will be the same as the method used for the bioburden determination. This testing will be conducted using test parameters outlined in ANSI/AAMI/ISO 11737-1.

7.3. Bioburden Determination

Ten samples each from three independent lots will be tested for bioburden using a specific test procedure. Each lot will have the results reported in colony forming units (CFU), with the ERF applied. The bioburden test will include aerobic bacteria and fungal enumerations.

7.4. Bioburden Test Method

The bioburden test method will be performed by immersion of the SIP. This testing will be conducted using test parameters defined in ANSI/AAMI/ISO 11737-1.

8. Verification Study Method

- 8.1. Verification Dose Determination
 - 8.1.1.A verification dose will be determined using the tables specified in AAMI TIR 33. The verification dose determination will be based on the final bioburden estimate for the product.
 - 8.1.2. Thirteen packaged verification samples will be sent to Sterigenics for irradiation at the calculated verification dose. The actual dose delivered will be measured and documented by Sterigenics. Sterigenics will send the samples directly to the contract laboratory after the verification dose has been performed.



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8.2. Verification Dose Range

If the maximum delivered dose exceeds the verification dose by more than 10%, the verification dose shall be repeated. If the maximum and minimum delivered dose is less than 90% of the verification dose, the verification dose may be repeated. If the delivered dose is less than 90% of the verification dose, and on the tests of sterility acceptable results are obtained, the verification experiment need not be repeated.

8.3. Test of Sterility

The test of sterility requires ten of the verification dose samples. The results will be reported as number of samples with growth and/or the number of samples with negative growth for the ten verification dose samples.

8.4. Test of Sterility Method

The test method will be performed by direct immersion. The test will be conducted using test parameters outlined in AAMI TIR 33 and ANSI/AAMI/ISO 11737-2.

8.5. Bacteriostasis/Fungistasis Determination

The bacteriostasis/fungistasis test requires three of the verification dose samples. This test is a validation of the test of sterility. This test will be conducted to verify that no bacteriostatic and fungistatic activity, which might compromise the sensitivity of the sterility test method, was exhibited by the samples, or formed by the verification dose exposure. The test results will be reported as a pass or fail.

8.6. Bacteriostasis/Fungistasis Test Method

The testing method will be the same as the method used for the test of sterility. The test will be performed using parameters outlined in the current USP.

9. Acceptance Criteria

Interpretation of results will be acceptable or unacceptable, as defined in AAMI TIR 33.

- 9.1. Verification is accepted if there are no more than one positive sterility test units out of the ten tested.
- 9.2. If two positives occur, perform another sterility test on ten additional units that have been irradiated at the same verification dose determined above. Upon completion of the second sterility test, add the number of positives from the two tests together. If the total number is two positives, verification is accepted.
- 9.3. If three or more positives occur, verification is not accepted. If the positive sterility units cannot be attributed to incorrect testing or dosing, an alternative dose determination method should be used unless the cause of the failure can be identified and corrected.
- 9.4. Bacteriostasis/Fungistasis test passes.



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10. Routine Sterilization

- 10.1. The routine sterilization dose is based on the acceptable results as outlined in Section 9 of this protocol. The 32.5-kGy minimum sterilization dose is considered valid for achieving a 10⁻⁶ SAL for the product listed.
- 10.2. With completion and acceptance of this validation study, subsequent batches of the product may be sterilized at a minimum dose of 32.5-kGy and released for use based only on the dosimeters indicating that the minimum SAL of 10-6 dose has been delivered to the product.
- 10.3. Dose audits must be performed according to an established schedule, as indicated in ANSI/AAMI/ISO 11137-1.

11. References

- 11.1.ANSI/AAMI/ISO 11737-1:2006, Sterilization of health care products Microbiological methods Part 1: Determination of the population of microorganisms on product
- 11.2.ANSI/AAMI/ISO 11737-2:2009, Sterilization of medical devices Microbiological methods Part 2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process
- 11.3.ANSI/AAMI/ISO 11137-1:2006, Sterilization of health care products Radiation Part 1: Requirements for development, validation, and routine control of a sterilization process for medical devices
- 11.4.ANSI/AAMI/ISO 11137-2:2006, Sterilization of health care products Radiation Part 2: Establishing the sterilization dose
- 11.5.AAMI TIR 33:2005, Sterilization of health care products Radiation Substantiation of a selected sterilization dose Method VD-Max
- 11.6.USP/NF, U.S. Pharmacopoeia (current version)



BIOBURDEN TEST REPORT

Procedure Reference: LB-MIC-004

SteriPro [®] Labs WO#	: 401711-MIC-004-I			Page	1 of 1
				Date Received	03/29/10
1400	erly-Clark Holcomb Bridge Road ell, GA 30076	Produ Cat / F Lot #:	Part #:	KIMTECH PURE* A5 (Apparel- Boots 2XL 12920 AR007702X	Cleanroom Sterile
Customer Specificat	tion Sheet# / Rev #:	I-004-411/ Rev.	2 Efficie	ency Recovery Factor	r (ERF): N/A
Test M Tryptic Soy Agar (TS Sabouraud Dextros Phosphate Buffered Sample Preparation	SA): e Agar (SDA): I Saline (PBS):	Manufacturer Biomerieux SteriPro SteriPro	IT ITC	Lot No. MFR432 01703132 02303143	Exp. Date 07/14/10 03/17/11 03/23/11
	Total Rinse Volume	e: 200mL	Volume Ex	tracted per Organisn	n Type:40mL
	Enumeration Metho		_	X Filtration _	Surface
	TSA Incubation Te	mperature: 30	-35°C SDA	A Incubation Tempera	ature: 20-25°C
Date Extracted/Incubated		Date TSA Enumerated:	04/02/1	Date SDA Enumerated:	04/05/10
_	TEST RESULTS (C	olony Forming U	nits (CFU) p		
	Sample ID		erobes (TSA)	Fungi-Yeast/Mold (SDA)	
	1		120	<5	li .
	2		130	<5 <5	
-	3 4		115	<5	
	5		135	10	
	6		235	5	
	7		150	10	
	8		135	5	
	9 10		175 125	5 <5	
	Total Average Bioburden/Devi	ce	144.0	6.0	
Cantral Beauties	TCA	0	SDA	0 PB \$	s 0
Control Results:	TSA				
Comments:			t	_	4/1/1
Prepared By:	- Min	<u> </u>		Date:	11/10
Reviewed By:	- Sugt	Masy		Date:	4/7/10

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BIOBURDEN TEST REPORT Procedure Reference: LB-MIC-004

SteriPro® Labs WO#:	401710-MIC-004-I		Page	1 of 1		
			Date Received:	03/29/10		
1400	erly-Clark Holcomb Bridge Road ell, GA 30076	Cat / Part #:	KIMTECH PURE* A5 C Apparel- Hood 88807 AR007502X	Cleanroom Sterile		
Customer Specification Sheet# / Rev #:I-004-411/ Rev. 2 Efficiency Recovery Factor (ERF):N/A						
Test Me Tryptic Soy Agar (TS Sabouraud Dextrose Phosphate Buffered Sample Preparation	Biome (SDA): Ster Ster Ster Ster Ster Ster Ster Ster	erieux IT iPro IT(iPro IT(Lot No. MFR432 01703132 02303143	Exp. Date 07/14/10 03/17/11 03/23/11		
Date Extracted/Incubated	Enumeration Method: TSA Incubation Temperatu Date TSA	Pour Plate ure: 30-35°C SDA uted: 04/02/		Surface ture: 20-25 ⁰ C		
	Sample ID	Aerobes (TSA)	Fungi-Yeast/Mold (SDA)			
-	1	175	<5			
<u> </u>	2	210	30			
-	3	200	5			
	4	180	5			
<u> </u>	5	175	5			
	6	200	<5			
Ì	7	215	10			
ļ	8	190	5			
Ī	9	270	15			
Ţ	10	125	10	i.		
	Total Average Bioburden/Device (calculated with ERF, if applicable)	194.0	9.5			
Control Results:	TSA 0	SDA	0 PBS	0		
Comments:	(10 1	a Am		11-10		
Prepared By:			Date:	7/11/		
Reviewed By:	- Bug Alla	it of the second se	Date:	4/7/10		

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BIOBURDEN EXHAUSTIVE RECOVERY VALIDATION TEST REPORT

Procedure Reference: LB-MIC-004

SteriPro® Labs V	NO#:	401712-MIC-004-I				Page	1_	of	2
						Date Recei	ved:	03	/29/10
1		-Clark comb Bridge Road GA 30076		duct: :/Part#: :#:				nroom	Sterile
Customer Speci	ification	Sheet# / Rev #:	I-004-411/ R	ev. 2	Efficiency F	Recovery Fa	ictor (El	RF):	N/A
<u>Te</u> Tryptic Soy Aga Sabouraud Dex Phosphate Buff	trose A	: gar (SDA):	Manufacture Biomerieux SteriPro SteriPro	<u>r</u>	Lot No ITMFR4 IT01703 IT02303	37 132	0	(p. Da 7/29/1 3/17/1 3/23/1	1
Sample Prepara	ation:	For each sample, ar	n SIP of 10% w	as cut and	d immersed	in PBS			
	ງ Page(od: Femperature: Date TSA Enumerated: Wash Results	Pour Plate 30-35°C		ed per Orga Filtration bation Tem Date SDA Enumera	peratur	_ Su e:	40mL rface 20-25°C
Comments: Prepared By: Reviewed By:	_	Alle	o Cus	ste		_ Date: _	4	Le Calle)

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BIOBURDEN EXHAUSTIVE RECOVERY VALIDATION TEST REPORT

Procedure Reference: LB-MIC-004

SteriPro [®] Labs WO#:	401712-MIC-004-I	Page	2	of	2

TEST RESULTS (Colony Forming Units (CFU) per Device):

FIRST WASH

Sample ID	Aerobes (TSA)	Fungi-Yeast/Mold (SDA)		
1	1850	50		
2	2200	150		
3	1050	50		
4	1850	<50		
5	1200	50		
Total Average Bioburden/Device	1630	70		

SECOND WASH

Sample ID	Aerobes (TSA)	Fungi-Yeast/Mold (SDA)
1	2300	50
2	1800	50
3	2850	50
4	2650	<50
5	1750	<50
Total Average Bioburden/Device	2270	50

THIRD WASH

Sample ID	Aerobes (TSA)	Fungi-Yeast/Mold (SDA)
1	2900	50
2	2150	100
3	2000	100
4	2550	50
5	1700	50
Total Average Bioburden/Device	2300	70

Control Results:	TSA	0	SDA	0	PBS	0

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BIOBURDEN TEST REPORT Procedure Reference: LB-MIC-004

	: 401709-MIC-004-I		Page	1 of1
			Date Recei	ved: 03/29/10
1400	erly-Clark Holcomb Bridge Road ell, GA 30076	Product: Cat / Part : Lot #:	Apparel- Coveralls	A5 Cleanroom Sterile -6XL
ustomer Specificat	tion Sheet# / Rev #:	I-004-411/ Rev. 2	Efficiency Recovery Fa	octor (ERF): 0.27
Test Me Fryptic Soy Agar (TS Sabouraud Dextrose Phosphate Buffered Sample Preparation	SA): e Agar (SDA): I Saline (PBS):	Manufacturer Biomerieux SteriPro SteriPro mple was cut and imm	<u>Lot No.</u> ITMFR432 IT01703132 IT02303143 nersed.	Exp. Date 07/14/10 03/17/11 03/23/11
	Total Rinse Volume	e:200mL Vo	lume Extracted per Orga	nism Type: 40mL
	Enumeration Metho	od: Pour Pla	ite X_ Filtration	Surface
	TSA Incubation Te	mperature: 30-35 ⁰	C SDA Incubation Tem	perature: 20-25°C
Date Extracted/Incubated	d: <u>03/30/10</u> E	Date TSA Enumerated:	04/02/10 Date SDA Enumera	
	TEST RESULTS (C	olony Forming Units	(CFU) per Device):	
Γ		A 1.		
l,	Sample ID	Aerob (TSA	1 –	old
-	Sample ID		(SDA)	old
-		(TSA	(SDA) (SDA)	old
-	1 2 3	(TSA 1750 1800 1850	(SDA) 0 <50 0 200 0 50	old
- - - -	1 2 3 4	(TSA 1750 1800 1850 3050	(SDA) (SDA)	old
-	1 2 3 4 5	(TSA 1750 1800 1850 3050 2250	(SDA) (SDA)	old
-	1 2 3 4 5 6	(TSA 1750 1800 1850 3050 2250	(SDA) (S	old
- - - - - - - -	1 2 3 4 5 6 7	(TSA 1750 1800 1850 3050 2250 1700 2950	(SDA) (S	old
-	1 2 3 4 5 6 7	(TSA 1750 1800 1850 3050 2250 1700 2950	(SDA) (S	old
-	1 2 3 4 5 6 7 8	(TSA 1750 1800 1850 3050 2250 1700 2950 2000	(SDA)	old
	1 2 3 4 5 6 7	(TSA 1750 1800 1850 3050 2250 1700 2950 2000 2750 1950	(SDA)	old
Control Results:	1 2 3 4 5 6 7 8 9 10 Total Average Bioburden/Devi (calculated with ERF, if ap	(TSA 1750 1800 1850 3050 2250 1700 2950 2000 2750 1950	(SDA)	PBS 0
Control Results:	1 2 3 4 5 6 7 8 9 10 Total Average Bioburden/Devi	(TSA 1750 1800 1850 3050 2250 1700 2950 2000 2750 1950 ce 8166	(SDA)	
Control Results: Comments: Prepared By:	1 2 3 4 5 6 7 8 9 10 Total Average Bioburden/Devi (calculated with ERF, if ap	(TSA 1750 1800 1850 3050 2250 1700 2950 2000 2750 1950 ce 8166	(SDA)	

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Procedure Reference: LB-MIC-004

Sterigenica International, Inc				
SteriPro® Labs WO#	: 407027-MIC-004-I		Page	e <u>1</u> of <u>1</u>
	-		Date Re	ceived: 04/07/10
1400	erly-Clark Holcomb Bridge Road vell, GA 30076	Product: Cat / Part Lot #:	Apparel-Covera	E* A5 Cleanroom Sterile alls-6XL
Customer Specifica	tion Sheet# / Rev #:	I-004-411/ Rev. 3	Efficiency Recovery	Factor (ERF): 0.27
Test M Tryptic Soy Agar (T Sabouraud Dextros Phosphate Buffered Sample Preparation	SA): se Agar (SDA): d Saline (PBS):	Manufacturer Biomerieux SteriPro SteriPro s cut and immersed i	Lot No. ITMFR437 IT01703132 IT00104165	Exp. Date 07/29/10 03/17/11 04/01/11
	Total Rinse Volume		lume Extracted per Or	
	Enumeration Metho			
	TSA Incubation Ter		C SDA Incubation To	
Date Extracted/Incubate		ate TSA numerated:	04/12/10 Date S Enum	DA erated: <u>04/13/10</u>
	TEST RESULTS (Co	olony Forming Units	(CFU) per Device):	
	Sample ID	Aerot (TS/		/Mold
	1	230		
	2	320		
	3	235		
	4	285		
	5	525		
	6 7	450 340		
	8	200	 	
	9	325		
	10	185		
	Total Average Bioburden/Devic	ce 1146		
Control Results:	TSA	0 SD	Δ0	PBS0
Comments: Prepared By:	door	Ocate) Date:	4/14/10
Droporod RV				\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
Prepared by.) (05M	<u></u> bato.	

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BIOBURDEN TEST REPORT

Procedure Reference: LB-MIC-004

	WO#: 407026-MIC-0	<u>04-l</u>		Page	
	, 			Date Received	: 04/07/10
ustomer:	Kimberly-Clark		Product:	KIMTECH PURE* A5	
	1400 Holcomb Bridge Ro	ad		Apparel-Coveralls-6XL	-
	Roswell, GA 30076		Cat / Part #:	12914 AR010402C	
			Lot #:	AR010402C	
Customer Spec	ification Sheet# / Rev #	:I-004-41	1/ Rev. 3 Effic	iency Recovery Facto	r (ERF): 0.27
Te	est Media	<u>Manufac</u>	turer	Lot No.	Exp. Date
Fryptic Soy Ag		Biomer	ieux	TMFR437	07/29/10
Sabouraud De	xtrose Agar (SDA):	SteriP		Г01703132	03/17/11
Phosphate But	fered Saline (PBS):	SteriF	Pro 1	Г00104165	04/01/11
Sample Prepar	ation: Each SIP of 10%	% was cut and	immersed in PBS.		
	Total Rinse Vo	lume: 20	00mL Volume E	Extracted per Organism	n Type: 40mL
	Enumeration N	lethod:	Pour Plate	X Filtration	Surface
	TSA Incubation	n Temperatur	e: 30-35 ⁰ C SI	A Incubation Tempera	ature: 20-25 ⁰ C
Date		Date TSA		Date SDA	
Extracted/Incu	bated:04/08/10	Enumerat	ed: <u>04/12</u>	2/10 Enumerated:	04/13/10
	Sample		rming Units (CFU)		_
	Campic	ID į	Aerobes (TSA)	Fungi-Yeast/Mold	
		ID	(TSA)	Fungi-Yeast/Mold (SDA) <50	
	11_	ID		(SDA)	
		ID	(TSA) 4250	(SDA) <50	
	1 2	ID	(TSA) 4250 4800	(SDA) <50 100 <50 100	
	1 2 3	ID	(TSA) 4250 4800 2050 2150 3900	(SDA) <50 100 <50 100 150	
	1 2 3 4 5 6	ID	4250 4800 2050 2150 3900 3100	(SDA) <50 100 <50 100 150 150	
	1 2 3 4 5 6	ID	(TSA) 4250 4800 2050 2150 3900 3100 1800	(SDA) <50 100 <50 100 150 150 <50	
	1 2 3 4 5 6 7	ID	(TSA) 4250 4800 2050 2150 3900 3100 1800 2250	(SDA) <50 100 <50 100 150 150 <50 50	
	1 2 3 4 5 6 7 8	ID	(TSA) 4250 4800 2050 2150 3900 3100 1800 2250 2700	(SDA) <50 100 <50 100 150 150 <50	
	1 2 3 4 5 6 7 8 9		(TSA) 4250 4800 2050 2150 3900 3100 1800 2250	(SDA) <50 100 <50 100 150 150 <50 50 50	
	1 2 3 4 5 6 7 8	rage Device	(TSA) 4250 4800 2050 2150 3900 3100 1800 2250 2700	(SDA) <50 100 <50 100 150 150 <50 50 50	
	1 2 3 4 5 6 7 8 9 10 Total Ave	rage Device	(TSA) 4250 4800 2050 2150 3900 3100 1800 2250 2700 2750	(SDA) <50 100 <50 100 150 150 <50 50 100	
Control Resu	1 2 3 4 5 6 7 8 9 10 Total Ave Bioburden/I (calculated with ERF	rage Device	(TSA) 4250 4800 2050 2150 3900 3100 1800 2250 2700 2750	(SDA) <50 100 <50 100 150 150 <50 50 100	s _ 0
Control Resu	1 2 3 4 5 6 7 8 9 10 Total Ave Bioburden/I (calculated with ERF	rage Device , if applicable)	(TSA) 4250 4800 2050 2150 3900 3100 1800 2250 2700 2750	(SDA) <50 100 <50 100 150 150 <50 50 50 50 100 314.8	s0
Comments:	1 2 3 4 5 6 7 8 9 10 Total Ave Bioburden/I (calculated with ERF	rage Device , if applicable)	(TSA) 4250 4800 2050 2150 3900 3100 1800 2250 2700 2750	(SDA) <50 100 <550 100 150 150 <50 50 50 314.8	s _ 0
	1 2 3 4 5 6 7 8 9 10 Total Ave Bioburden/I (calculated with ERF	rage Device , if applicable)	(TSA) 4250 4800 2050 2150 3900 3100 1800 2250 2700 2750	(SDA) <50 100 <50 100 150 150 <50 50 50 50 100 314.8	s
Comments:	1 2 3 4 5 6 7 8 9 10 Total Ave Bioburden/I (calculated with ERF	rage Device , if applicable)	(TSA) 4250 4800 2050 2150 3900 3100 1800 2250 2700 2750	(SDA) <50 100 <550 100 150 150 <50 50 50 314.8	s _ 0

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Dose Calculation for Kimberly-Clark KIMTECH PURE*A5 Cleanroom Sterile Apparel

Based on the Total Recoverable Aerobic and Fungi Bioburden results from three independent lots of the KIMTECH PURE* A5 Cleanroom Sterile Apparel, the theoretical overall average taken from the three lots was selected to calculate the subprocess Verification dose. The Recovery Factor was determined to be 0.27.

Per AAMI TIR33: Method VD-Max30 guidelines, the Sub-process Verification dose is based on the overall average bioburden from three lots, unless one of the averages is two or more times greater than that of that overall average. The verification dose was derived from the theoretical bioburden average of the three lots. The theoretical lot averages are as follows:

Total Bioburden (Aerobic and Fungal)

Lot Number

As colony forming units (cfu) per device

AR007602A 8666.7 cfu/device

AR010402B 11777.8 cfu/device

AR010402C 11333.3 cfu/device

The overall average bioburden for the three lots, including the recovery factor, is 10592.6 cfu per device. Based on 10592.6 cfu per device, and utilizing the SIP verification dose calculation, the verification dose was determined directly from Table A.7 of the AAMI TIR33 guidance. The verification dose for the device is **9.9 kGy**.

In accordance with the AAMI TIR33: Method VD-Max30 guidelines, ten units of product should be sampled from a single production lot and irradiated at the verification dose of 9.9 kGy \pm 10%. After irradiation the samples will be sent to SteriPro Labs to be placed on test of sterility.

If these ten units, which will have been irradiated at 9.9 kGy, meet the criteria set forth in the AAMI TIR33: Method VD-Max30 guidelines (no more than one positive per ten units), then the minimum sterilization dose from a sterility assurance level would be 30.0 kGy.

Best regards,

Zabrina Tumaitis

Consultant, SteriPro Consulting



Certificate of Processing

STERIGENICS 344 Bonnie Circle Corona CA 92880 TEL 951 340-0700 FAX www.sterigenics.com

R55480102 RIS0003

16:05:51 GMT 04/19/10

Page - 1 of 1

Customer Name: SteriPro Consulting

P.O.#

12334270 / KIMBERLY CLARK

Processing Facility: Corona

Work Order #

412952

Sales Order #

363808 **GMT**

X_STERIPRO_18

Irradiation Date/Time:

04/18/10

17:03:00

EXCELL

Customer Item Number /

Customer Item Description

Customer Lot Number /

Customer Load Number

UOM 101.00 CA

450.00

X_CARTON_STERIPRO_18

AR007602B COVERALLS - 6XL

CA Total

Quality Test Summary

------Signed By -----

Op# Quality Test Description 450.00 Minimum Dose

Pass/Fail User Result Pass JGARCIA 9.4 kGy

Irradiation Cell:

Date 1 Time 04/19/10 01:38:28 GMT

Maximum Dose

Reason Code Test

Pass JGARCIA

Jose Garcia

10.2 kGy

Jose Garcia

Reason Code Test

04/19/10 01:38:44 GMT

Sterigenics certifies that the materials listed above (as described by the Manufacturer) received the indicated doses within the precision and accuracy of

the dosimetry system employed.

Electronically Signed By:

Annette Posada

Work Order Completions

Date: 04/19/10

16:00:47 GMT



STERILITY VALIDATION (B&F) TEST REPORT Procedure Reference: LB-MIC-027

SteriPro® La	bs WO#: 415045-MIC	C-027-I		Pa	age <u>1</u>	of <u>1</u>		
Date Receiv	ed: <u>04/20/10</u>	Date On-Tes	st: 04/21/10	_ Date O	ff-Test: <u>0</u>	4/26/10		
Customer:	Kimberly Clark 1400 Holcomb Bridge Roswell, GA 30076	Road	Product: Cat / Part #: Lot #: Load #: Sterilization Run Date Sterilized:	Sterile A 12914 AR0076 N/A	opparel- Co	A5 Cleanroor veralls -6XL	m	
1	est Media	Manufactu	ırer <u>Volum</u>		<u>ot No.</u> 1204185	Exp. Date 04/12/11		
	TSB	SteriPro	3000m		1904185	04/12/11		
Product Ste	rility Customer Specif	ication Sheeta	# / Revision:	I-036-797/F	Rev. 0			
	Test Method: Immersion of SIP							
Test Results	<u>X</u> P	ASS	FA	IIL				
				Т	SB			
	ORGANISM		Cor	itrol		ample		
Bacillus subt	ilis (ATCC #6633)			+		+		
Candida albic	cans (ATCC #10231)			+		+		
Aspergillus n	iger (ATCC #16404)			+		+		
Comments: N/A								
Prepared By	r Jen	Nota	aught-	Date:	4/2	1/10		
Reviewed B	y:			Date:	4/2	7/10		
		1"				•		

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PRODUCT STERILITY TEST REPORT

Procedure Reference: LB-MIC-036

SteriPro® Labs WO#:	415039-MIC-036-I		Page1	of <u>1</u>
Date Received: 04/	20/10 Date On-Te	est: 04/21/10	Date Off-Test: _	05/05/10
Customer: Kimberly - 1400 Holo Roswell, C	omb Bridge Road	Product: Cat / Part #: Lot #: Load #: Sterilization Run: Date Sterilized:	KIMTECH PURE* A Sterile Appareal-Co 12914 AR007602B AR007602B 412952 04/19/10	
<u>Test Media</u>	<u>Manufact</u>	turer <u>Volume</u>	Lot No.	Exp. Date
Tryptic Soy Broth ((TSB) SteriPro	Labs 3000mL	IT01204185	04/12/11
Number of Samples:	10			
Customer Specificatio	n Sheet# / Revision #:	I-036-797/ Rev. 0		
Test Method: Immersion of SIP.				
Test Results: TSB: Open Control: Results Meet Criteria:	X No Growth X No Growth X YES	 S NO	_ Growth _ Growth	
Attachments Included	: N/A			
<u>Comments</u> :	N/A			
Prepared By:	Gathern	str	Date:	50
Reviewed By:	Catherin	Reat	Date: <i>5</i> /	6/10

The test results relate only to the samples as provided and tested. This report may not be reproduced, except in full, without written approval from SteriPro Labs.

Dose setting Report

Kimberly-Clark Corporation

1400 Holcomb Bridge Road, Roswell, Georgia

The following sample(s) was/were submitted and identified on behalf of the applicant as:

Kimtech Pure A5 Sterile Integrated Hood and Mask

Date of Sample Receipt

2013/03/04

Lab Test ID

13-033A

Date on Test

2013/03/04

Test Item

Dose Setting

Test Method

ISO 11137-2-2012: Method 1

Signature

Fight She 2013-04-01
Edited by Date

Inke Gang 2013 04-01
Reviewed by Date

Spagle Xm Jois-04-01 Sheream Mc Cory 203-04-01

Approved by Date Kimberly-Clork Professional

Revision

0	2013/04/01	First edition
Revision No.	Date	Revision Description

Document N°:

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Revision N°: 1

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Tested sample description

Sample information				
Sample name Kimtech Pure A5 Sterile Integrated Hood and Mask				
Product code	36072			
Lot#	First batch, second batch, third batch			
Number of Sample	138 samples			
SIP	1.0			

No.: DS004-13r

Sample photo



Test results

Bioburden validation (Inoculation)

Sample ID	- Counted	PF	Recovered per sample(CFU)	Confirmed Inoculation level(CFU)	ERF
1	14		28		0.56
2	13		26		0.52
3	14	2.0	28	52/48	0.56
4	13		26	26	0.52
5	10		20		0.40
Average	12.8	NA	25.6	50	0.51
C.F (Corre	ection Factor)			2.0	

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♦ Bioburden test

	Bior	ourden Test	Result (Samp	e Lot: First	batch)		
ID	CFU count		CFU	P.F	C.F	CFU/ SIP sample	
	Aerobes	Fungi	Counted		0.1	Or Or Oil Sampl	
1	164	87	251			2008	
2	135	22	157			1256	
3	221	87	308			2464	
4	156	112	268			2144	
5	142	47	189		0.0	1512	
6	348	80	428	4	2.0	3424	
7	98	45	143			1144	
8	125	8	133			1064	
9	147	48	195			1560	
10	127	64	191			1528	
control	0	0	N/A	N/A	N/A	N/A	
SIP Average			1810.4 C	FU/SIP sam	ole		
Batch 1 Average			1810.4	CFU/Device		and the state of t	
	Biobu	ırden Test R	esult (Sample	Lot: Second	d batch)		
ID.	CFU count		CFU	D.E.	O.F.	CELI/SID somn	
ID	Aerobes	Fungi	Counted	P.F	C.F	CFU/ SIP sample	
1	259	67	326			2608	
2	91	40	131			1048	
3	301	68	369			2952	
4	424	42	466			3728	
5	147	52	199	4	2.0	1592	
6	324	124	448			3584	
7	235	68	303			2424	
8	184	10	194			1552	

No.: DS004-13r

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	sterioenics -	
7	Sterigenics	

9	242	62	304			2432		
10	187	74	261			2088		
control	0	0	N/A	N/A	N/A	N/A		
SIP Average	2400.8 CFU/SIP sample							
Batch 2 Average	2400.8 CFU/Device							
	Biok	ourden Test	Result (Samp	le Lot: Third	batch)			
Ī.	CFU	count	CFU	D.F.	0.5	OF WORD		
ID	Aerobes	Fungi	Counted	P.F	C.F	CFU/ SIP sample		
1	145	56	201			1608		
2	149	10	159			1272		
3	260	56	316			2528		
4	145	48	193			1544		
5	140	34	174	4	2.0	1392		
6	256	52	308	4	2.0	2464		
7	88	79	167			1336		
8	98	25	123			984		
9	231	60	291			2328		
10	296	84	380			3040		
control	0	0	N/A	N/A	N/A	N/A		
SIP Average			1849.6 (CFU/SIP sam	ple			
Batch 3 Average			1849.6	CFU/Device				

♦ Establishing sterilization dose

Establishing sterilization dose						
Overall average bioburden(SIP)	Verification dose (SIP sample)	Overall average bioburden (Device)	Sterilization dose (10 ⁻⁶)			
2020.3 CFU	12.0 kGy	2020.3 CFU	26.1 kGy			

Note: Overall average bioburden= (batch1+batch2+batch3)/3= (1810.4+2400.8+1849.6)/3= 2020.3 CFU/device

♦ Irradiation of verification dose

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No.: DS004-13r

Verification dose is irradiated on 100 samples (Lot: XN306301X) within the ±10% of target dose 12.0 kGy. Detail information refers to **Certificate of Irradiation for Test Sample** from **Sterigenics Shanghai E-Beam Co., Ltd (PO20130314128391)**.

Sterility test validation (Bacteriostasis & Fungistasis Test)

Sterility Test Validation (B/F) Result						
Culture	Bacillus subtilis ATCC 6633	Candida albicans ATCC10231	Aspergillus brasiliensis ATCC16404			
Test Sample	Positive	Positive	Positive			
Inoculated Control	Positive	Positive	Positive			
Inoculum Level(CFU)	31	39	30			
Conclusion		Pure A5 Sterile Integrated riostatic and fungistatic properties or ough the test.				

♦ Sterility test

Sterility Test Result					
SIP	1.0				
Sample number	100				
Type of Media	TSB				
Media Volume	800mL				
Incubation Period	14 days				
Incubation Temperature	28°C - 32°C				
Results	1 Positive				

Conclusion

From validation results show, the product of **Kimtech Pure A5 Sterile Integrated Hood and Mask (Code: 36072) PASS** the dose setting by using Method 1 according to ISO 11137-2. A sterility assurance level (SAL) of 10⁻⁶ is confirmed when irradiating the product by a sterilization dose of 26.1 kGy for routine production.

References

- ♦ ISO 11737-1:2006 Sterilization of health care products Microbiological methods Part 1: Determination of the population of microorganisms on product
- ISO 11737-2:2009 Sterilization of medical devices—Microbiological methods---Part2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process

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♦ ISO 11137-2:2006 Sterilization of health care products---Radiation---Part 2: Establishing the sterilization dose

No.: DS004-13r

- ♦ Sterigenics internal Work Instruction SHEB-WI-LB-MIC-003
- ♦ Sterigenics internal Work Instruction SHEB-WI-LB-MIC-016

Remarks

- ♦ The test report is invalid unless signed by editor, reviewer and approver.
- ♦ The test report is invalid if altered.
- ♦ Please contact testing laboratory within 15 days after receiving the test report if any objection to the report, otherwise it will not be accepted.
- ♦ The test report is only responsible for the sample provided by customer.
- ♦ The test report cannot be used for any commercial purpose unless approved by testing laboratory.
- ♦ Tested by SteriPro laboratory, Sterigenics Shanghai E-Beam Ltd. No.588 ChuanTu Road, Chuansha. Pudong, Shanghai, China. 201202. Tel(86-21) 58594680 Fax(86-21)58599310

=========End of report===============

Dose setting Report

Kimberly-Clark Corporation

1400 Holcomb Bridge Road, Roswell, Georgia

The following sample(s) was/were submitted and identified on behalf of the applicant as:

Kimtech Pure A5 Sterile Sleeves

Date of Sample Receipt

2013/03/04

Lab Test ID

13-033A

Date on Test

2013/03/04

Test Item

Dose Setting

Test Method

ISO 11137-2-2012: Method 1

Signature

Reviewed by

2013-04-01 Spagle Xu Approved by

Thursa MCCon 2013-04-61 Kimberly-Clark Frofessional

Revision

0	2013/04/01	First edition	
Revision No.	Date	Revision Description	

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Tested sample description

Sample information				
Sample name	Kimtech Pure A5 Sterile Sleeves			
Product code	36077			
Lot [#]	First batch, second batch, third batch			
Number of Sample	138samples			
SIP	1.0			

No.: DS005-13r

Sample photo



Test results

Bioburden validation (Inoculation)

	Bioburden validation test results(Sample Lot: First batch)						
Sample ID	Counted	PF	Recovered per sample(CFU)	Confirmed Inoculation level(CFU)	ERF		
1	22		33.0		0.66		
2	18		27.0		0.54		
3	25	1.5	37.5	52/48	0.75		
4	16		24.0		0.48		
5	21		31.5		0.63		
Average	20.4	NA	30.6	50	0.61		
C.F (Corre	ection Factor)			1.7			

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♦ Bioburden test

	District of the second		Result (Sampl	e Lot. First	Daterry			
ID	CFU count		CFU	P.F	C.F	CFU/ SIP sample		
	Aerobes	Fungi	Counted		0.1	Or or on sample		
1	1620	147	1767	3		9011.7		
2	2240	259	2499			12744.9		
3	2410	84	2494			12719.4		
4	2520	138	2658			13555.8		
5	1780	187	1967			10031.7		
6	740	124	864		1.7	4406.4		
7	960	67	1027			5237.7		
8	1780	252	2032			10363.2		
9	2520	51	2571			13112.1		
10	1460	54	1514			7721.4		
control	0	0	N/A	N/A	N/A	N/A		
SIP Average			9890.4 C	FU/SIP sam	ole			
Batch 1 Average			9890.4	CFU/Device)			
	Biobu	ırden Test R	esult (Sample	Lot: Second	d batch)			
ID	CFU count		CFU			OF IVE		
	Aerobes	Fungi	Counted	P.F	C.F	CFU/ SIP sample		
1	1180	50	1230	3	1.7	6273		
2	117	26	143			729.3		
3	222	47	269			1371.9		
4	680	32	712			3631.2		
5	146	55	201			1025.1		
6	210	51	261			1331.1		
7	226	56	282					1438.2
8	123	85	208			1060.8		

No.: DS005-13r

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Sterigenics

No.: DS005-13r

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9	112	38	150			765
10	292	82	374			1907.4
control	0	0	N/A	N/A	N/A	N/A
SIP Average	1953.3 CFU/SIP sample					
Batch 2 Average	1953.3 CFU/Device					
	Biob	urden Test	Result (Samp	le Lot: Third	batch)	
ID	CFU d	count	CFU	P.F	C.F	CFU/ SIP sample
	Aerobes	Fungi	Counted			
1	1460	152	1612	3		8221.2
2	2100	39	2139			10908.9
3	1760	56	1816			9261.6
4	980	56	1036			5283.6
5	1540	82	1622		1.7	8272.2
6	420	21	441			2249.1
7	680	95	775			3952.5
8	1440	65	1505			7675.5
9	1980	34	2014			10271.4
10	1460	127	1587			8093.7
control	0	0	N/A	N/A	N/A	N/A
SIP Average	7419.0 CFU/SIP sample					
Batch 3 Average			7419.0	CFU/Device		

Establishing sterilization dose

Establishing sterilization dose				
Overall average bioburden(SIP)	Verification dose (SIP sample)	Overall average bioburden (Device)	Sterilization dose (10 ⁻⁶)	
6420.9 CFU	13.6 Gy	6420.9 CFU	27.9 KGy	

Note: Overall average bioburden= (batch1+batch2+batch3)/3= (9890.4+1953.3+7419.0)/3= 6420.9 CFU/device

Irradiation of verification dose

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Verification dose is irradiated on 100 samples (Lot: XN306301X) within the ±10% of target dose 13.6 kGy. Detail information refers to Certificate of Irradiation for Test Sample from Sterigenics Shanghai E-Beam Co., Ltd (PO20130313128358).

No.: DS005-13r

Sterility test validation (Bacteriostasis & Fungistasis Test)

Sterility Test Validation (B/F) Result					
Culture	Bacillus subtilis ATCC 6633	Candida albicans ATCC10231	Aspergillus brasiliensis ATCC16404		
Test Sample	Positive	Positive	Positive		
Inoculated Control	Positive	Positive	Positive		
Inoculum Level(CFU)	31	39	30		
Conclusion	The products of Kimtech Pure A5 Sterile Sleeves (Code: 36077) do not exist bacteriostatic and fungistatic properties, or such properties have been eliminated through the test.				

Sterility test

Sterility Test Result			
SIP	1.0		
Sample number	100		
Type of Media	TSB		
Media Volume	400mL		
Incubation Period	14 days		
Incubation Temperature	28°C - 32°C		
Results	2 Positive		

Conclusion

From validation results show, the product of Kimtech Pure A5 Sterile Sleeves (Code: 36077) PASS the dose setting by using Method 1 according to ISO 11137-2. A sterility assurance level (SAL) of 10⁻⁶ is confirmed when irradiating the product by a sterilization dose of 27.9 kGy for routine production.

References

- ♦ ISO 11737-1:2006 Sterilization of health care products Microbiological methods Part 1: Determination of the population of microorganisms on product
- ♦ ISO 11737-2:2009 Sterilization of medical devices—Microbiological methods—Part2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process

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No.: DS005-13r

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======End of report=============

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